

SENATE BILL No. 201

DIGEST OF SB 201 (Updated January 31, 2007 1:37 pm - DI 104)

Citations Affected: IC 12-15.

Synopsis: Medicaid pharmacy survey and preferred drug list report. Changes the timing from twice per year to one time per year for the drug utilization review board report concerning the preferred drug list for Medicaid recipients. Repeals provisions requiring the office of Medicaid policy and planning to conduct a survey of pharmacy providers to assess the appropriate level of pharmacy dispensing fees.

Effective: July 1, 2007.

Miller

January 8, 2007, read first time and referred to Committee on Health and Provider Services. February 1, 2007, amended, reported favorably — Do Pass.





First Regular Session 115th General Assembly (2007)

PRINTING CODE. Amendments: Whenever an existing statute (or a section of the Indiana Constitution) is being amended, the text of the existing provision will appear in this style type, additions will appear in this style type, and deletions will appear in this style type.

Additions: Whenever a new statutory provision is being enacted (or a new constitutional provision adopted), the text of the new provision will appear in **this style type**. Also, the word **NEW** will appear in that style type in the introductory clause of each SECTION that adds a new provision to the Indiana Code or the Indiana Constitution.

Conflict reconciliation: Text in a statute in *this style type* or *this style type* reconciles conflicts between statutes enacted by the 2006 Regular Session of the General Assembly.

C

SENATE BILL No. 201

O

A BILL FOR AN ACT to amend the Indiana Code concerning Medicaid.

p

Be it enacted by the General Assembly of the State of Indiana:

y

- SECTION 1. IC 12-15-35-28, AS AMENDED BY P.L.101-2005, SECTION 3, IS AMENDED TO READ AS FOLLOWS [EFFECTIVE JULY 1, 2007]: Sec. 28. (a) The board has the following duties:
 - (1) The adoption of rules to carry out this chapter, in accordance with the provisions of IC 4-22-2 and subject to any office approval that is required by the federal Omnibus Budget Reconciliation Act of 1990 under Public Law 101-508 and its implementing regulations.
 - (2) The implementation of a Medicaid retrospective and prospective DUR program as outlined in this chapter, including the approval of software programs to be used by the pharmacist for prospective DUR and recommendations concerning the provisions of the contractual agreement between the state and any other entity that will be processing and reviewing Medicaid drug claims and profiles for the DUR program under this chapter.
 - (3) The development and application of the predetermined criteria and standards for appropriate prescribing to be used in



1

2

3

4

5

6

7

8

9

10

11 12

13

14

1516

17

1	retrospective and prospective DUR to ensure that such criteria
2	and standards for appropriate prescribing are based on the
3	compendia and developed with professional input with provisions
4	for timely revisions and assessments as necessary.
5	(4) The development, selection, application, and assessment of
6	interventions for physicians, pharmacists, and patients that are
7	educational and not punitive in nature.
8	(5) The publication of an annual report that must be subject to
9	public comment before issuance to the federal Department of
10	Health and Human Services and to the Indiana legislative council
11	by December 1 of each year. The report issued to the legislative
12	council must be in an electronic format under IC 5-14-6.
13	(6) The development of a working agreement for the board to
14	clarify the areas of responsibility with related boards or agencies,
15	including the following:
16	(A) The Indiana board of pharmacy.
17	(B) The medical licensing board of Indiana.
18	(C) The SURS staff.
19	(7) The establishment of a grievance and appeals process for
20	physicians or pharmacists under this chapter.
21	(8) The publication and dissemination of educational information
22	to physicians and pharmacists regarding the board and the DUR
23	program, including information on the following:
24	(A) Identifying and reducing the frequency of patterns of
25	fraud, abuse, gross overuse, or inappropriate or medically
26	unnecessary care among physicians, pharmacists, and
27	recipients.
28	(B) Potential or actual severe or adverse reactions to drugs.
29	(C) Therapeutic appropriateness.
30	(D) Overutilization or underutilization.
31	(E) Appropriate use of generic drugs.
32	(F) Therapeutic duplication.
33	(G) Drug-disease contraindications.
34	(H) Drug-drug interactions.
35	(I) Incorrect drug dosage and duration of drug treatment.
36	(J) Drug allergy interactions.
37	(K) Clinical abuse and misuse.
38	(9) The adoption and implementation of procedures designed to
39	ensure the confidentiality of any information collected, stored,
40	retrieved, assessed, or analyzed by the board, staff to the board, or
41	contractors to the DUR program that identifies individual



42

physicians, pharmacists, or recipients.

1	(10) The implementation of additional drug utilization review
2	with respect to drugs dispensed to residents of nursing facilities
3	shall not be required if the nursing facility is in compliance with
4	the drug regimen procedures under 410 IAC 16.2-3.1 and 42 CFR
5	483.60.
6	(11) The research, development, and approval of a preferred drug
7	list for:
8	(A) Medicaid's fee for service program;
9	(B) Medicaid's primary care case management program;
10	(C) Medicaid's risk based managed care program, if the office
11	provides a prescription drug benefit and subject to IC 12-15-5;
12	and
13	(D) the children's health insurance program under IC 12-17.6;
14	in consultation with the therapeutics committee.
15	(12) The approval of the review and maintenance of the preferred
16	drug list at least two (2) times per year.
17	(13) The preparation and submission of a report concerning the
18	preferred drug list at least two (2) times one (1) time per year to
19	the select joint commission on Medicaid oversight established by
20	IC 2-5-26-3.
21	(14) The collection of data reflecting prescribing patterns related
22	to treatment of children diagnosed with attention deficit disorder
23	or attention deficit hyperactivity disorder.
24	(15) Advising the Indiana comprehensive health insurance
25	association established by IC 27-8-10-2.1 concerning
26	implementation of chronic disease management and
27	pharmaceutical management programs under IC 27-8-10-3.5.
28	(b) The board shall use the clinical expertise of the therapeutics
29	committee in developing a preferred drug list. The board shall also
30	consider expert testimony in the development of a preferred drug list.
31	(c) In researching and developing a preferred drug list under
32	subsection (a)(11), the board shall do the following:
33	(1) Use literature abstracting technology.
34	(2) Use commonly accepted guidance principles of disease
35	management.
36	(3) Develop therapeutic classifications for the preferred drug list.
37	(4) Give primary consideration to the clinical efficacy or
38	appropriateness of a particular drug in treating a specific medical
39	condition.
40	(5) Include in any cost effectiveness considerations the cost
41	implications of other components of the state's Medicaid program



42

and other state funded programs.

1	(d) Prior authorization is required for coverage under a program
2	described in subsection (a)(11) of a drug that is not included on the
3	preferred drug list.
4	(e) The board shall determine whether to include a single source
5	covered outpatient drug that is newly approved by the federal Food and
6	Drug Administration on the preferred drug list not later than sixty (60)
7	days after the date on which the manufacturer notifies the board in
8	writing of the drug's approval. However, if the board determines that
9	there is inadequate information about the drug available to the board
10	to make a determination, the board may have an additional sixty (60)
11	days to make a determination from the date that the board receives
12	adequate information to perform the board's review. Prior authorization
13	may not be automatically required for a single source drug that is newly
14	approved by the federal Food and Drug Administration, and that is:
15	(1) in a therapeutic classification:
16	(A) that has not been reviewed by the board; and
17	(B) for which prior authorization is not required; or
18	(2) the sole drug in a new therapeutic classification that has not
19	been reviewed by the board.
20	(f) The board may not exclude a drug from the preferred drug list
21	based solely on price.
22	(g) The following requirements apply to a preferred drug list
23	developed under subsection (a)(11):
24	(1) Except as provided by IC 12-15-35.5-3(b) and
25	IC 12-15-35.5-3(c), the office or the board may require prior
26	authorization for a drug that is included on the preferred drug list
27	under the following circumstances:
28	(A) To override a prospective drug utilization review alert.
29	(B) To permit reimbursement for a medically necessary brand
30	name drug that is subject to generic substitution under
31	IC 16-42-22-10.
32	(C) To prevent fraud, abuse, waste, overutilization, or
33	inappropriate utilization.
34	(D) To permit implementation of a disease management
35	program.
36	(E) To implement other initiatives permitted by state or federal
37	law.
38	(2) All drugs described in IC 12-15-35.5-3(b) must be included on
39	the preferred drug list.
40	(3) The office may add a drug that has been approved by the
41	federal Food and Drug Administration to the preferred drug list



42

without prior approval from the board.

1	(4) The board may add a drug that has been approved by the	
2	federal Food and Drug Administration to the preferred drug list.	
3	(h) At least two (2) times one (1) time each year, the board shall	
4	provide a report to the select joint commission on Medicaid oversight	
5	established by IC 2-5-26-3. The report must contain the following	
6	information:	
7	(1) The cost of administering the preferred drug list.	
8	(2) Any increase in Medicaid physician, laboratory, or hospital	
9	costs or in other state funded programs as a result of the preferred	
10	drug list.	
11	(3) The impact of the preferred drug list on the ability of a	
12	Medicaid recipient to obtain prescription drugs.	
13	(4) The number of times prior authorization was requested, and	
14	the number of times prior authorization was:	
15	(A) approved; and	_
16	(B) disapproved.	
17	(i) The board shall provide the first report required under subsection	
18	(h) not later than six (6) months after the board submits an initial	
19	preferred drug list to the office.	
20	SECTION 2. IC 12-15-31.1 IS REPEALED [EFFECTIVE JULY 1,	
21	2007].	
		_



COMMITTEE REPORT

Madam President: The Senate Committee on Health and Provider Services, to which was referred Senate Bill No. 201, has had the same under consideration and begs leave to report the same back to the Senate with the recommendation that said bill be AMENDED as follows:

Page 3, reset in roman line 17.

Page 3, line 18, reset in roman "preferred drug list at least".

Page 3, line 18, after "times" insert "one (1) time".

Page 3, line 18, reset in roman "per year to the select joint".

Page 3, reset in roman line 19.

Page 3, line 20, reset in roman "(14)".

Page 3, line 20, delete "(13)".

Page 3, line 23, reset in roman "(15)".

Page 3, line 23, delete "(14)".

Page 5, line 2, reset in roman "(h) At least".

Page 5, line 2, after "times" insert "one (1) time".

Page 5, line 2, reset in roman "each year, the board shall provide a report".

Page 5, reset in roman lines 3 through 14.

and when so amended that said bill do pass.

(Reference is to SB 201 as introduced.)

MILLER, Chairperson

Committee Vote: Yeas 9, Nays 1.

y

